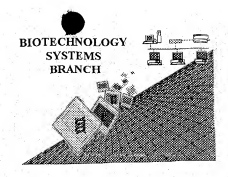
RAW SEQUENCE LISTING ERROR REPORT



The Biotechnology Systems Branch of the Scientific and Technical Information Center (STIC) detected errors when processing the following computer readable form:

Application Serial Number:	09/761,142	
Source:	OIPE	
Date Processed by STIC:	1/30/20	00/

THE ATTACHED PRINTOUT EXPLAINS DETECTED ERRORS.
PLEASE FORWARD THIS INFORMATION TO THE APPLICANT BY EITHER:

1) INCLUDING A COPY OF THIS PRINTOUT IN YOUR NEXT COMMUNICATION TO THE APPLICANT, WITH A NOTICE TO COMPLY or,

2) TELEPHONING APPLICANT AND FAXING A COPY OF THIS PRINTOUT, WITH A NOTICE TO COMPLY FOR CRF SUBMISSION QUESTIONS, PLEASE CONTACT MARK SPENCER, 703-308-4212.

FOR SEQUENCE RULES INTERPRETATION, PLEASE CONTACT ROBERT WAX, 703-308-4216. PATENTIN 2.1 e-mail help: patin21help@uspto.gov or phone 703-306-4119 (R. Wax) PATENTIN 3.0 e-mail help: patin3help@uspto.gov or phone 703-306-4119 (R. Wax)

TO REDUCE ERRORED SEQUENCE LISTINGS, PLEASE USE THE <u>CHECKER</u> <u>VERSION 3.0 PROGRAM</u>, ACCESSIBLE THROUGH THE U.S. PATENT AND TRADEMARK OFFICE WEBSITE. SEE BELOW:

Checker Version 3.0

The Checker Version 3.0 application is a state-of the-art Windows based software program employing a logical and intuitive user-interface to check whether a sequence listing is in compliance with format and content rules. Checker Version 3.0 works for sequence listings generated for the original version of 37 CFR §§1.821 – 1.825 effective October 1, 1990 (old rules) and the revised version (new rules) effective July 1, 1998 as well as World Intellectual Property Organization (WIPO) Standard ST.25.

Checker Version 3.0 replaces the previous DOS-based version of Checker, and is Y2K-compliant. Checker allows public users to check sequence listings in Computer Readable form (CRF) before submitting them to the United States Patent and Trademark Office (USPTO). Use of Checker prior to filing the sequence listing is expected to result in fewer errored sequence listings, thus saving time and money.

Checker Version 3.0 can be down loaded from the USPTO website at the following address: http://www.uspto.gov/web/offices/pac/checker

VERIFICATION SUMMARY

PATENT APPLICATION: US/09/761,142 TIME: 11:19:35

DATE: 01/30/2001

Input Set : A:\31133A.ST25.txt

Output Set: N:\CRF3\01302001\1761142.raw

```
L:13 M:270 C: Current Application Number differs, Replaced Current Application No
L:13 M:271 C: Current Filing Date differs, Replaced Current Filing Date
L:1221 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:26
L:1221 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:26 L:1221 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:26
L:1221 M:258 W: Mandatory Feature missing, <223> not found for SEQ 1D#:26 L:1221 M:340 W: (46) "n" or "Xaa" used: Feature required, for SEQ 1D#:26
\text{L}:1285~\text{M}:258~\text{W}: Mandatory Feature missing, <220> not found for SEQ ID#:26
L:1285 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:26
\text{L}:1285\ \text{M}:258\ \text{W}: Mandatory Teature missing, <222> not found for SEQ TD#:26
L:1285 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:26
M:340 Repeated in SeqNo=26
L:1286 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:26
L:1286 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:26
L:1286 M:258 W: Mandatory Feature missing, <222> not found for SEQ 1D#:26
L:1286 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:26
L:1287 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:26
L:1287 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:26 L:1287 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:26
L:1287 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:26
L:1839 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:32
L:1839 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:32 L:1839 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:32
L:1839 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:32
L:1839 M:340 W: (46) "n" or "Xaa" used: Feature required, for SEQ ID#:32
L:11047 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:153
L:11047 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:153 L:11047 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:153
L:11047 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:153 L:11047 M:340 W: (46) "n" or "Xaa" used: Feature required, for SEQ ID#:153
L:19618 M:258 W: Mandatory Feature missing, <220> not found for SEQ TD#:258
L:19618 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:258
L:19618 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:258
L:19618 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:258
L:19618 M:340 W: (46) "n" or "Xaa" used: Feature required, for SEQ ID#:258
L:28080 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:356
L:28080 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:356
L:28080 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:356
L:28080 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:356
L:28080 M:340 W: (46) "n" or "Xaa" used: Feature required, for SEQ ID#:356
```

```
<210> 26
<211> 6413
<212> DNA
<213> Drosophila melanogaster
<400> 26
gaatteette acaggeggtt tettggeege attategtta atattgttgt tggacatgta 60
cagctcgcat tgctgcacta ttttatccaa ttttcgaatt tcgctcagtt ccacgcaaga 120
tttgtgggac atttttaagt ctgcaaagta tagaaacgaa aaagtgttgc aaattataat 180
caaacattcg caaagctcag aatatccgga ttcgtaagca aattcaaata aatcttcgct 240
taaaaacaac ttaataataa atccaaaatt tccacttact ccagattgaa attggcgaaa 300
actgcgattt cttacctaat gcactttctt atttacttgc aaatcaaaac agaacaggta 360
tgacatgcac gcggagaaat caggagtgga aaatcatttt caatcaatta aaaacacttc 420
aattogatot tgttogttac acatactota otttttaaat atcatagata aaataaaaaa 480
atgaatagat gtatgtaaat acatggcatt ttcatttcga agtcaactgc tttataagac 540
caactttttt ttcccggtgt aatcaggctt tgcttttctg aaattttcag acatgcgtac 600
tgctttcggt tttcaaagat tcgaaaccac cgtgattttc gaaccattgt tattttcctt 660
ctcgcattcg gtcacactat tttcaaacgt aaacaattga tttttcctgc taaacaaaag 720
tggttgaaaa tgtcgggaat cgctgccaaa aagattgccg aagcggagga cctggtgaag 780
caggoogaga agaggtatoo ocaacaaaaa tatoottata aaggggttto caaataacac 840
ccatctttat ttgcagcttg aagttgtcca tgctgaaatg ggttcctgat tacgatagtg 900
ctgcggatga gtattccaaa gctggtgagt ttcacaaagt aggagtgact aatatcaccc 960
agagtateta tetataaaaa cattgtgaca teaatgagat atgtaaatae tatgaattae 1020
agctaacgat tatcccaaag tatccataac ttgcaagata cacacatatt ggtgtacaat 1080
tatgtatett tettttaeat ttttaattge agecaetgea tategaatag etaaaagtta 1140
tgataagagc aaggagtgtt ttctgaaggc aatcgacgcc tataaaaaca acaagtcttg 1200
qttccatqct qcaaaqqcat acqaacaqqt qaqcttaaat tgaagttaat ataagtgaag 1260
ttaataggaa atttaaaccc tctaaatgtg cacatettet agateatttt gctgtcaaag 1320
gatgeegata agetacaega agetgaggaa tacgeeaaca aateggegag tttgtateaa 1380
cagcacggtt ccccagaggc agccgcatcc gccttggata aagccgccaa gttaactgaa 1440
tecaageate etgacatgge titgegette tateageatg etetagaagt cataatggtt 1500
cgtcctattt gacactgctc tagtcaattc cattcttaaa ctattaattt tatacttacc 1560
cgacagatcg aggattccgt ccgtcaagca gccgagtatg atcaaaagtt tccaggatac 1620
tggtcaaact aaggaggtat gtacctatga ttgcaacgaa tcactttggg tatccttagc 1680
ttatatcatc caatcactcc aggtacgacg aagccacgaa tgcgctcaaa aaggagatca 1740
gcttgaatca gcaaacggaa tcttacggac aaattggacg cctagttgtg gccttggtga 1800
tggtccaact ggctcgcggg gattccgtgg aagccgaaaa gaccttcagg gagtggggaa 1860
agtacgetat tetatetaaa aacatteeae taceteaggg tatacagatg gagaagaagg 2040
etggegacae tgetgetgtg agtatteta teetaaaaat ggaattegaa attetegtt 2100

(Partial bitus of Seguene 26 - due to Ena Sunnay
sige of sequene, only first ena shown - Heet
similar enars exist in the rest of the
sequence)
```

FJI

Please Note:

Use of n and/or Xaa have been detected in the Sequence Listing. Please review the Sequence Listing to ensure that a corresponding explanation is presented in the <220> to <223> fields of each sequence which presents at least one n or Xaa.

RAW SEQUENCE LISTING

DATE: 01/30/2001 PATENT APPLICATION: US/09/761,142 TIME: 11:19:34

Input Set : A:\31133A.ST25.txt

Output Set: N:\CRF3\01302001\I761142.raw

```
275 atttaccggg cgttaagacc caccgaatgc ggttaattta agtgataata taatatta 420
276 aataagaata tgttaatacg tgtaccetgt gacaaatgtt tacagtotot aactaatteg 480
277 tactttegta tittacaaga acgtaaacgt acgatatcag agtatctaaa agtaaaacca 540
278 egaateytte atgtttaegg tittaeggatt aegagtaaaa tggtateaaa tettitgate 600
279 aactacctat cgaccattac agactgacgg tttacattca acttttttac tgcgacatct 660
280 gegattaett ageaegaaca taaataaata aacegtgggg tegtgatgtg caegttaaat 720
281 autoauttte acquacaqeq ecqtaatttt atautaatee agteatgttt tacatataaa 780
282 totgaaactt ogacogacta gttgotaaat tattoattga aagtggataa titttattat 840
283 ttgggatttt gataatgegg aatageeate caeaatttat tattaegtaa ggtatatega 900
284 tcaaggactt caagtatgcg tgttggaacc gltcattaaa taattcaata accccggtgt 960
285 gaaaatcaaa atcaaagatt gagacgetet ttetaeaage aagttettet ttatetaate 1020
286 etteattama teaggtatta aaaggaatea tgatteetga attgattgat aaaacamagg 1080
287 gaaagcaaqt etgeaegttg cacaagaaga agaggacace geaetggtgg acceeaagee 1140
288 ggcagaggca gatgcogteg atacceagce gtgaaaegce teetegteaa aageeteage 1200
289 ygaaggaccc gtacatgcac cggaacttcc acagcatgtt cataaggtal atgcactcat 1260
290 ttgacgcaag tgttgacgtt cagtgtctga ttggagaaac gtaagtcatg gattaaacgt 1320
291 caccggacca atateggegg gacccgagga tggagaagtg gaagtgcege taegeceteg 1380
292 tegactacce gtecaagete etegegtacg egetggagaa getegettte tgggtgtege 1440
293 tgctgttcta cgtcggtcac gtgtcggāga agccgacgac gccgtaactc cctggcgtcc 1500
294 tgatqqaaqt cetectegLq eegegggacq ggtcgtcgac gacgatgegg aagetgacga 1560
295 ggtteggeeg egtgeagatg etecteecta egaggtggtt eegecatega tgagaetett 1620
296 aegteegeet egagitgatg atgtegaegt aeegetagta aegggagete eaetetgtaa 1680
297 titiggtatgc gtcccattit aagtgaacgg tgttcccatg aactcgtgtt cccatacaaa 1740
298 geogtgoage gaatacette teaalaggtt altgaaaact taattgegtg tgttgactga 1800
299 gtcacttgtg taaaattgat gaaaaaataa gcagtgcgaa tttatcttta gggacggttc 1860
300 tattetacae acaaaaaagt aagtttggta aaaacaatga tttcaaaaga aatttgattt 1920
301 atacagtitic caegotagea atgataaata catgigicaa caactegaaa aegaactaca 1980
302 atatticaat atticatcat tectgeaatt etgteatitt ataaaaaaat tgitteaage 2040
303 qaaqaataa gotaaaqota aggtatagtt ataatataat aatgttaaat acataaaaag 2100
304 toaaaaaccc gaacaagtgg cgaatggtaa acccgtteeg agegtteata eggttttggt 2160
305 totaltteet geteetetgg gggtagttge tgact
308 <210> SEQ ID NO: 17
309 <21.1> LENGTH: 227
310 <212> TYPE: PRT
311 <213> ORGANISM: Drosophila melanogaster
313 <400> SEQUENCE: 17
314 Met Glu Lys Ser Phe Pro Ile Thr Pro Trp Lys Tyr Gly Leu Leu Val
31.5 1 5
317 Thr Cys ile Leu ile Val Thr Cys Asn Val Phe Phe Ser Cys Gly 318 20 25 30
320 Val Thr Thr Trp Gly Ser Ala Val Ser Val Tyr Gly Ser Tyr Gly Ser 321 \phantom{-}35\phantom{+}40\phantom{+}45\phantom{+}
323 Ala Leu Cys Gly Gly Ala Val Phe Gly Val Ala Phe Leu Gly Met Tyr
324 50 55 60
326 Val Ala Leu Lys Val Ser Tyr Lys Tyr Ser Ile Tyr Tyr Leu Ile Cys
327 65 70 75 80
329 Ser Gly Leu Val Tle Ala Ala Leu Gly Ser Tyr Leu Phe Thr Phe Thr
330 85 90 95
332 Ala Met Arg Glu Gln Leu Met Gly Arg Phe Glu Glu Arg Met Arg Asp
```

RAW SEQUENCE LISTING PATENT APPLICATION: US/09/761,142

DATE: 01/30/2001 TIME: 11:19:34

Input Set : A:\31133A.ST25.txt

Output Set: N:\CRF3\01302001\I761142.raw

```
205 acgeggalet geageagace ecegaecace tetalelgga egggtgeate gaaaaggtge 660
206 agagetteta egaaagegae aagetgeget teateatagt gteetgggtg etagtggeet 720
207 togagttaat etgettegee tiggeegigt tietggeeat tagtittaag aacaageage 780
208 gacggatgga gttetagtte taggettegg taatetegag etatecaaca gtacaaacte 840
209 ggaategggg teteget.gat attittetet teaacattte ataaccaaat geaaaggaca 900
210 gtoataaatt attoactoot accttaatgt aacetgtaat taaagtacat atttgtagtt 960
211 caattaccca ttataagtat cataataaat gtgcgcgtgt ttgttttcac atga
214 <210> SEQ ID NO: 15
215 <211> LENGTH: 208
216 <212> TYPE: PRT
217 <213> ORGANISM: Drosophila melanogaster
219 <400> SEQUENCE: 15
220 Met Gly Cys Ala Thr Thr Ser Val Lys Ile Ala Ser Ile Val Leu Asn
                                          1.0
223 Ala Val Leu Gly Phe Leu Ala Ala Gly Ala Ile Gly Trp Ile Ala Tyr
224 20 25 30
226 Asn Ala Asp Thr Glu Thr Glu Glu Phe Val Ile Ala Ala Tyr Ile Ala
229 Cys Ser Leu Ile Leu Val Phe Ala Leu Leu Gly Ile Phe Ala Ala Ile
       5.0
                             55
232 Arg Glu Ser Val Val Leu Thr Ala Thr Ser Ala Val Phe Leu Leu Ile 233 \, 65 \, 70 \, 75 \, 80
235 Leu Ala Ile Leu Gln Ile Val Ser Thr Cys Leu Phe Leu His Glu Phe
                     85
                                          90
238 Asp Val Lys Ser Gly Arg Asp Met Val Glu Val Ala Trp Gln Ala Asn
                                     1.05
239
               100
241 Asn Met Asp Ser Leu Gln Gln Lys His Glu Cys Cys Gly Gln Ser Ser 242 115 120 125
244 Ala Gln Asp Tyr Ile His Leu Ser Leu Leu Ile Pro Pro Ser Cys Tyr 245 \phantom{-}130\phantom{0} 135 \phantom{-}140\phantom{0}
247 Ala Asp Leu Gln Gin Thr Pro Asp His Leu Tyr Leu Asp Gly Cys Ile
248 145
             150
                                            155
250 Glu Lys Val Gln Ser Phe Tyr Glu Ser Asp Lys Leu Arg Phe Ile Ile
                 1.65
                                      170
                                                            175
253 Val Ser Trp Val Leu Val Ala Phe Glu Leu Ile Cys Phe Ala Leu Ala
254 180 185
256 Val Phe Leu Ala Ile Ser Phe Lys Asn Lys Gln Arg Arg Met Glu Phe
257 195
                                 200
                                                      205
263 <210> SEQ TD NO: 16
264 <211> LENGTH: 2195
265 <212> TYPE: DNA
266 <213> ORGANISM: Drosophila melanogaster
268 <400> SEQUENCE: 16
269 tacctictica gealaggita ttgtgggace tteatgeegg atgaceagtg gaegtaagag 60
270 taacaccatt cactttttct acgogtaaag gggtaaggto togaaagggt aaaagggoto 120
271 ggactgggtt taccettaga aegtgttgtg ggecaetaeg etegttaete ggttttaget 180
272 tittagitat aacgogggit caagigtaga aagggttote tegitittaa ceactgitae 240
273 teggegattt etgatatttg tttagetaeg ttagagtttt ttataaatat ttaegtatta 300
274 tggacaagtg tatgaatgta ttaaggegge eggtegacaa acattaaaaa egttatagat 360
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RAW SEQUENCE LISTING PATENT APPLICATION: US/09/761,142

DATE: 01/30/2001 TIME: 11:19:34

Input Set : A:\31133A.ST25.txt

Output Set: N:\CRF3\01302001\1761142.raw

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139 <212> TYPE: DNA
140 <213> ORGANISM: Artificial Sequence
142 <220> FEATURE:
143 <223> OTHER INFORMATION: Description of Artificial Sequence: PCR Primer
144
         Sequence Sp3
146 <400> SEQUENCE: 10
                                                                      24
147 gagtacgcaa agetttaact aigt
150 <210> SEQ ID NO: 11
151 <211> LENGTH: 23
152 <212> TYPE: DNA
153 <213> ORGANISM: Artificial Sequence
155 <220> FEATURE:
156 <223> OTHER INFORMATION: Description of Artificial Sequence: PCR Primer
157
         Sequence Sp6
159 <400> SEQUENCE: 11
160 tgaccacate caaacatect ett
                                                                      2.3
163 <210> SEQ ID NO: 12
164 <211> LENGTH: 25
165 <212> TYPE: DNA
166 <213> ORGANISM: Artificial Sequence
168 <220> FEATURE:
169 <223> OTHER INFORMATION: Description of Artificial Sequence: PCR Primer
170
         Sequence Sp5
172 <400> SEQUENCE: 12
173 gcatcacaaa aatcgacget caagt
176 <210> SEQ ID NO: 13
177 <211> LENGTH: 19
178 <212> TYPE: DNA
179 <213> ORGANISM: Artificial Sequence
181 <220> FEATURE:
182 <223> OTHER INFORMATION: Description of Artificial Sequence: PCR Primer
1.83
         Sequence Spep1
185 <400> SEQUENCE: 13
                                                                      19
186 gacactcaga atactattc
189 <21.0> SEQ ID NO: 14
190 <233> LENGTH: 1014
191 <212> TYPE: DNA
192 <213> ORGANISM: Drosophila melanogaster
194 <400> SEQUENCE: 14
195 agogaacgca gaacgcgcca gaccaaaaag ttcagattcg agagcggata tcccggcgag 60
196 cgttcacqqa aatatatttg tttgttattc gagtccagca acgaatattt aaataaacaa 120
197 aaaacqaact ttattogtgt goggagagag aagtcaaaag alccaataaa atgggttgcg 180
198 ccacqaccaq cqtgaagatc gootccatcg ttotgaatgc cgttttaggg ttottgctg 240
199 ctggggccat cggctggata gcttacaatg cggacacgga gacggaggaa ttcgtaatag 300
200 ccgcttacat cgcgtgeteg ctcatcetgg tetttgetet gctgggcatc ttcgcggcca 360
201 teegggaate ggtggtgetg aetgeaacga gtgetgtett eetgetgate ttggeeatee 420
202 tgcagatcgt gagcacctgc ctgttcctcc acgagttcga cgtgaagagc ggccgggaca 480
203 tggtggaggt ggcetggcag gcgaacaaca tggatteett gcagcagaag cacgagtget 540
204 geggecagag cagegeccag gactatatec accteageet gelgateceg eccagetget 600
```

RAW SEQUENCE LISTING
PATENT APPLICATION: US/09/761,142

DATE: 01/30/2001 TIME: 11:19:34

Input Set : A:\31133A.ST25.txt
Output Set: N:\CRF3\01302001\1761142.raw

68 <400> SEQUENCE: 4	
69 ccttagcatg tccgtggggt ttgaat	26
72 <210> SEQ ID NO: 5	
73 <21.1> LENGTH: 28	
74 <212> TYPE: DNA	
75 <213> ORGANISM: Artificial Sequence	
77 <220> FEATURE:	
78 <223> OTHER INFORMATION: Description of Artificial Sequence	e: PCR Primer
79 Sequence Pry2	
81 <400> SEQUENCE: 5	
82 cttgccgacg ggaccacctt atgttatt	28
85 <210> SEQ ID NO: 6	
86 <211> LENGTH: 19	
87 <212> TYPE: DNA	
88 <213> ORGANISM: Artificial Sequence	
90 <220> FEATURE:	
91 <223> OTHER INFORMATION: Description of Artificial Sequence	e: PCR Primer
92 Sequence Plw3-1	
94 <400> SEQUENCE: 6	
95 tgtcggcgtc atcaactcc	19
98 <210> SEQ ID NO: 7	
99 <211> LENGTH: 29	
100 <212> TYPE: DNA	
101 <213> ORGANISM: Artificial Sequence	
103 <220> FEATURE:	
104 <223> OTHER INFORMATION: Description of Artificial Sequen	ce: PCR Primer
1.05 Sequence Pwht1	
107 <400> SEQUENCE: 7	
108 gtaacgctaa tcactccgaa caggtcaca	29
111 <210> SEQ ID NO: 8	
112 <211> LENGTH: 25	
113 <212> TYPE: DNA	
114 <213> ORGANISM: Artificial Sequence	
116 <220> FEATURE:	
117 <223> OTHER INFORMATION: Description of Artificial Sequen	ce: PCR Primer
118 Sequence Splac2	
120 <400> SEQUENCE: 8	
121 gaatteactg geogtegttt tacaa	25
124 <210> SEQ ID NO: 9	
1.25 <21.1> LENGTH: 22	
126 <212> TYPE: DNA	
127 <213> ORGANISM: Artificial Sequence	
129 <220> FEATURE:	
130 <223> OTHER INFORMATION: Description of Artificial Sequen	ce. PCR Primar
131 Sequence Spl	OC. FOR FIRMSI
133 <400> SEQUENCE: 9	22
134 acacacctt tecteteaac aa	44
137 <210> SEQ ID NO: 10	
138 <211> LENGTH: 24	

Raw Sequence Listing Error Summary

	ERROR DETECTED	SUGGESTED CORRECTION SERIAL NUMBER: UT/ 10/11
A TTN	. NEW DITLES CASES. I	LEASE DISREGARD ENGLISH "ALPHA" HEADERS, WHICH WERE INSERTED BY PTO SOFTWARE
ALIN		The number/text at the end of each line "wrapped" down to the next line.
1	Wrapped Nucleics	This may occur if your file was retrieved in a word processor after creating it.
		Please adjust your right margin to .3, as this will prevent "wrapping".
2	Wrapped Aminos	The amino acid number/text at the end of each line "wrapped" down to the next line.
	• • • • • • • • • • • • • • • • • • •	This may occur if your file was retrieved in a word processor after creating it.
		Please adjust your right margin to .3, as this will prevent "wrapping".
		r lease adjust your right margin to .o., do this tim protont the party.
3	Incorrect Line Length	The rules require that a line not exceed 72 characters in length. This includes spaces.
·	. Wiles	
4	Misaligned Amino Acid	The numbering under each 5th amino acid is misaligned. This may be caused by the use of tabs
	Numbering	between the numbering. It is recommended to delete any tabs and use spacing between the numbers.
	Numbering	between the manipening. It is recommended to the second se
5	Non-ASCII	This file was not saved in ASCII (DOS) text, as required by the Sequence Rules.
٠ <u> </u>	11011710011	Please ensure your subsequent submission is saved in ASCII text so that it can be processed.
		Tiedase cristice your subsequent additional to a service and a service a
6	Variable Length	Sequence(s) contain n's or Xaa's which represented more than one residue.
·	7 2.1.2.1.5 2.1.1.g	As per the rules, each n or Xaa can only represent a single residue.
	-	Please present the maximum number of each residue having variable length and
		indicate in the (ix) feature section that some may be missing.
		indicate in the (ix) readure section that some may be massing.
7	Patentin ver. 2.0 "bug"	A "bug" in PatentIn version 2.0 has caused the <220>-<223> section to be missing from amino acid
′	Fateritin ver. 2.0 bug	sequence(s) Normally, PatentIn would automatically generate this section from the
		previously coded nucleic acid sequence. Please manually copy the relevant <220>-<223> section
		to the subsequent amino acid sequence. This applies primarily to the mandatory <220>-<223>
		sections for Artificial or Unknown sequences.
_		the street state was the following format for each plinned accurage:
8	Skipped Sequences	Sequence(s) missing. If intentional, please use the following format for each skipped sequence:
	(OLD RULES)	(2) INFORMATION FOR SEQ ID NO:X:
		(i) SEQUENCE CHARACTERISTICS:(Do not insert any headings under "SEQUENCE CHARACTERISTICS")
		(xi) SEQUENCE DESCRIPTION:SEQ ID NO:X:
		This sequence is intentionally skipped
		Please also adjust the "(iii) NUMBER OF SEQUENCES:" response to include the skipped sequence(s).
9	Skipped Sequences	Sequence(s) missing. If intentional, please use the following format for each skipped sequence.
2.	(NEW RULES)	<210> sequence id number
- 1		₹400> sequence id number
1		000
	Use of n's or Xaa's	Use of n's and/or Xaa's have been detected in the Sequence Listing.
	(NEW RULES)"	Use of <220> to <223> is MANDATORY if n's or Xaa's are present.
		In <220> to <223> section, please explain location of n or Xaa, and which residue n or Xaa represents.
1	Use of <213>Organism	Sequence(s) are missing this mandatory field or its response.
	(NEW RULES)	
2	Use of <220>Feature	Sequence(s) are missing the <220>Feature and associated headings.
0	(NEW RULES)	Use of <220> to <223> is MANDATORY if <213>ORGANISM is "Artificial" or "Unknown"
		Please explain source of genetic material in <220> to <223> section.
	•	(See "Federal Register," 6/01/98, Vol. 63, No. 104, pp. 29631-32) (Sec. 1.823 of new Rules
		(000 1 600 at 1.020 a
,	Det # 0.0 # - "	Please do not use "Copy to Disk" function of Patentln version 2.0. This causes a corrupted
· ——	Patentin ver, 2.0 "bug"	
	The same of the sa	file, Testitling in missing mandatory numeric identifiers and responses (as indicated on raw sequence listing).
		Instead, please use "File Manager" or any other means to copy file to floppy disk.

OIPE

RAW SEQUENCE LISTING

PATENT APPLICATION: US/09/761,142

DATE: 01/30/2001 TIME: 11:19:34

. Input Set : A:\31133A.ST25,txt

Output Set: N:\CRF3\01302001\1761142.raw

Does Not Comply Corrected Diskette Needed

3 <110> APPLICANT: Kamdar, Kim Broadus, Julie Stam, Lynn Bachmann, Jane 8 <120> TITLE OF INVENTION: NUCLEIC ACID SEQUENCES FROM DROSOPHILA MELANOGASTER THAT ENCODE PROTEINS ESSENTIAL FOR LARVAL VIABILITY AND USES THEREOF 11 <130> FILE REFERENCE: PB/5-31133A 13 <140> CURRENT APPLICATION NUMBER: US/09/761,142 13 <141> CURRENT FILING DATE: 2001-01-16 13 <150> PRIOR APPLICATION NUMBER: US 60/176,418 14 <151> PRIOR FILING DATE: 2000-01-14 16 <160> NUMBER OF SEQ ID NOS: 361 18 <170> SOFTWARE: PatentIn Ver. 2.1 20 <210> SEQ ID NO: 1 21 <211> LENGTH: 27 22 <212> TYPE: DNA 23 <213> ORGANISM: Artificial Sequence 25 <220> FEATURE: 26 <223> OTHER INFORMATION: Description of Artificial Sequence: PCR Primer 27 Sequence Plac4 29 <400> SEQUENCE: 1 30 actglgcgtt aggtcctgtt cattgtt 27 33 <210> SEQ ID NO: 2 34 <211> LENGTH: 24 35 <212> TYPE: DNA 36 <213> ORGANISM: Artificial Sequence 38 <220> FEATURE: 39 <223> OTHER INFORMATION: Description of Artificial Sequence: PCR Primer Sequence Plac1 42 <400> SEQUENCE: 2 43 cacccaagge tetgeteeca caat 24 46 <210> SEQ ID NO: 3 47 <211> LENGTH: 23 48 <212> TYPE: DNA 49 <213> ORGANISM: Artificial Sequence 51 <220> FEATURE: 52 <223> OTHER INFORMATION: Description of Artificial Sequence: PCR Primer 53 Sequence Pry4 55 <400> SEQUENCE: 3 56 caateatate get.gtctcae tea 23 59 <210> SEQ ID NO: 4 60 <211> LENGTH: 26 61 <212> TYPE: DNA 62 <213> ORGANISM: Artificial Sequence 64 <220> FEATURE: 65 <223> OTHER INFORMATION: Description of Artificial Sequence: PCR Primer